4153261467

) hereby certify that this correspondence is being deposited with the United States Postal S first class moil in an envelope addressed to: Had Doked ORIGINAL Washington, D.C. 20231, on 12/8/95

Attorney Docket No. 016994-003122

IN THE UNITED STATES STONE OF THE STATE OF THE

In the applicant part of

H MHORE, ET AL.

mediant No. 1 optimize the

Filed: November 16, 1993

For: PRODUCTION OF RECOMBINANT POLYPEPTIDES BY BOVINE SPECIES AND TRANSGENIC

METHODS

Examiner, J. Grammers.

Art Unit: 1804

DECLARATION OF NEAL FIRST UNDER 37 CFR 5132

Commissioner of Patents and Trademarks Awalington, D.C. 20201

3111

- 1, Real J., First, State as follows
- 1. My present position is Professor of Reproductive Biology and Animal Biotechnology at the University of Wisconsin. I am a consultant for several companies including Pharming μ ν , the assignee of the above-captioned application (the '015 application). A copy of my curriculum wither is attached . I have been asked by Pharming B.V. to give my opinion of the claims to methods of generating transgenic bovines in the '019 application in view of the commerce in the office action mailed May 5, 1995.
- 2. In forming my opinion, I have reviewed the total application, the office action mailed May 2, 1995, and pertinent references cited therein. One of these references if First, US 5 231,979 (the 1979 patent), of which I am the first-named historia or
- 3. I note that the 1019 application is directed (in part) to methods of producing transgenic bovines (hereafter the Pharming methods). The methods involve the following steps: harvesting immature occytos from hovinos, culturing the immature occytes in vitro, fertilizing the occytes in vitro to produce



H. DOBOER, ET AL. Serial No.: 08/154,019 Page 2

PATENT

zygotes, introducing a transgene into the zygotes, culturing the zygotes in vitro to form a embryo and transplanting the embryo into a female bovine. The application also exemplifies the successful use of the methods to produce a transgenic bovine.

- 4. I understand that the Examiner takes the position that the successful application of the Pharming methods to achieve a transgenic bovine would have been obvious from the '979 patent. I respectfully disagree with this position for the reasons stated below. To the contrary, I believe that the successful practice of the Pharming methods represents a substantial advance in the art whose attainment was not reasonably expected from, e.g. the '979 patent.
- 5. The Pharming methods represent a substantial advance in the art because quite surprisingly they made possible a y der range of genetic manipulations than that performed to the 'org patent. The '979 patent discusses the development of an in virro culturing method that allowed production of viable boving blastoysts. The '979 patent of First et al. is directed to generic methods for the culture and co-culture of boving employs.

The DeBoer et al. '019 application is not directed to a method of culturing bovine embryos per se, but rather uses the method in development of a system for producing transgenic cattle. It is the system and its successful application that is novel and inventive for cattle in the DeBoer application

Our goal underlying the method in the 1979 patent was the manipulation of blastocyst-stage embryos (as described in column 1, lines 55-60). For example, this procedure would expedite the generation of herds of genetically superior animals (particularly dairy cattle) by allowing cloning or blastocyst believed to have desirable naturally occurring characteristics. We did not perform any manipulations involving introduction of transgenes into embryonic cells. By contrast, the Pharming methods allow introduction of any transgene into a zygote leading to phenotypes

H. DOBOER, ET AL. Serial No.: 08/154,019 Page 3

HUV UT 1000 17:44

١

PATENT

not found in nature, such as a cow producing milk containing a human protein.

At the time of tiling the '97' patent we did know that transgenesis was possible in made and sheep and we inought, might some day be possible in cattle. Thus, in the '979 patent, we noted that in vitro culture might be used in genetic engineering (column 1, line 62). However, we did not describe particular transgenesis procedures (e.g., embryonic stem cells, microinjection of zygotes or infection with retroviruses) or indicate how our in vitro culture method might be adapted to the exploited in any of these procedures.

- 6. In may opinion, the successful practice of the Pharming methods could not reasonably have been expected from the method described in the 1979 patent. This opinion is based in part, on the following facts.
- (a) The scientific literature as of December 1989 (the effective filing date of the above captioned application) indicated that attempts to produce transgenic bovines up to then had proved extremely difficult, lengthy and expensive. For example, one review article reports:

Most scientists working on transgenic animals—he it for improving traits such as feed efficiency or for using them as factories for human pharmaceuticals—shy away from cattle. In the cow...you have a three-four year project. And it's a costly venture as well

Van Brunt, Bio/Technology 6, 1149-1154 (1988) at p 1152
As of Docomber 1989, there were no confirmed reports in a pagereviewed scientific journal of any viable transgenic bovine calf
having been produced. The Biery, Lostkutoff and Bondioli
references mentioned in the office action discuss only attempts
which did not generate transgenic bovine calves. Although a very
low frequency of expression was obtained in early fetuses. Today
expression in embryos and early fetuses is known to often be from
non chromosome integrated DNA (Krisher et al., Animal
Biotechnology 6, 15-25 (1995); Bowen et al., Biol. Reprod. 50,
664-668 (1994)). In light of this general background of failure

1

H. DOBOER, ET AL. Serial No.: 08/154,019 Page 4

PATENT

and frustration, most practitioners (including me) would have approached alternate well-ode or generation transgent confiderable a measure of skepticism and an expectation that considerable empirical experimentation lay ahead.

- (b) There would have been a number of problems and uncertainties in trying to dombine the in vitro procedure with the poorly successful bovine methods of Biery, Lostkutoff and Bondioli, in which occyte maturation and fertilization are performed in vivo, but transgenic offspring did not result For example, it was unpredictable whether traditional microinse tion procedures for in-vivo eggs could have been successfully applied without modification to in-vitro matured pocytes matured oocytes might have different physiological properties (e.g., structural differences in the zona pellucida (the propertyTayer surrounding the occyte, its hardening etc) due to the different environment in which maturation accurred; to my knowledge, this had never been thoroughly investigated in any species. It was also unpredictable whether the phasing of the cell cycle of in-vitro occytes would have been different or impact on the visibility of the pronucleus and therefore on the injection protocol. Difficulty may also have related to the relative timing of microninjection and fertilization which a unit not be the same for in-vivo and in-vitro matured cocytes (1) ... relative timing would have been expected to be important in development of a successful protocol, because if the DNA was injected before S phase, it might be degraded before interpation and after S phase, it could not be integrated until the tro will Chattery ...
- (c) A further source of unpredictability was whether the block on bovine embryo development in vitro occurring at about the 8-cell stage could be overcome, and if so, with what officiency, in the context of a transgenesis protocol. The tare patent describes how the block on bovine embryo development could be overcome at an efficiency of about 20% by supplementing culture media with epithelial cells in the context of the aloring protocol used. This was an empirical observation; at that time, we did not understand the mechanism by which the block occurred

H. DOBOER, ET AL. Serial No.: 08/154,019 Page 5

PATENT

or by which it was selleved. There in on it having onlying no a manipulation (i.e., indeceduate them) that on it have a consideral impact on cell physiology to a phasing of the cell cycle) it was unclear, whether and to what extent epithelial cell suppresentation of collumn medium would be effective in consideral the cell blockage at the eight-cell stage. Today there are totally defined media culture systems effective in culturing boying embryos to blastocyst (i.e., Rosenkrane & First, it Animal, Sci. 72, 434-437 (1994)). This was not considered possible when the '979 patent studies were done.

The first of the same of the same of

uncertainties discussed in the previous paragraph might know required considerable symmatrical to perfect or observed vet the ultimate end noise for the efficient of such experimentation (i.e., the production of a viable transger bovine) would not have been apparent until several years of so that practically it was not possible to vary systematically most of the parameters. These factors explain my view that the provision and demonstrated officacy of Pharming's transcomment methods were not a routine development.

8. That others hold similar views regarding the substantial and dramatic advance of the Pharming's method to a difficult and unfruitful field is illustrated by the following comments

The commercial development of transgenic bovine technologies, however, has been frustrated between the protocols used successfully with smaller animals—which require large numbers of embryon and several surgical procedures—are prohibitively expensive when applied to cattle. The establishment of an in vitro embryo production system, as described by Herman de Boer and him coworkers at Gene Pharming Europe...is therefore a dramatic breakthrough in chlarging the transgenic pharm—yard.

whole, rade balances quart

Į į

A group in the Netherlands reports in a paper in the September issue of Bio/Technology successful generation of the first transports dating call.

II. 🌠 BALU ALTO lest Available Copy

H. DOBOER, ET AL.

9: 1000 1::40

PATENT

Serial No.: 08/154,019 Page 6

> which carries a gene for production in committeet human la televete (sid) - combern esse un a re-a novel fetegrares en viet grove e nor génerale p transgenic cattle.

Selzer, Chemical & Engineering News 69, 7 (1991)

But historically, efforts to produce transgen. dairy cows have been thwarted because of cumbersome and costly surgical procedures. however, researchers from Gene Pharming Europe. have discumvented the need for surgical removal and transfer of embryos by combining gone transwith an in vitro embryo production system.

Finally, I wish to state for the record that the co. . repose of this declaration is to assist in assessment or th parentability of the Pharming claims, and the validity and accuraif the claims in the 1979 patent involve different issues

I have been duly warned that willfu! false statements and tie like are publishable by fine and imprisonment or both under Section 100; of little is of the United States Code and that willful false statements may jeopardize the validity of the above-identified patent application or any parent results

Meal & Wird

1 6 30 -

lebas 1 - First

Nov. 8 1995

Dated